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METHOD OF USING ZOT OR ZONULIN TO INHIBIT LYMPHOCYTE PROLIFERATION IN AN ANTIGEN-SPECIFIC MANNER

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CROSS-REFERENCE TO RELATED APPLICATIONS

This application is an application filed under 35 U.S.C. § 111(a) claiming benefit pursuant to 35 U.S.C. § 119(e)(i) of the filing date of the Provisional Application No. 60/100,266, filed September 14, 1998, pursuant to 35 U.S.C. § 111(b).

15 <u>FIELD OF THE INVENTION</u>

The present invention relates to antigen-specific down-regulation of an immune response using Zot or zonulin. Specifically, the present invention provides a method for inhibiting antigen presenting cell-mediated antigen-specific lymphocyte proliferation in a dose-dependent manner by administering an effective amount of Zot or zonulin.

25 <u>BACKGROUND OF THE INVENTION</u>

I. Tight Junctions and the Actin Cytoskeleton

The tight junctions (hereinafter "tj") or zonula occludens (hereinafter "ZO") are one of the hallmarks of absorptive and secretory epithelia (Madara, J. Clin. Invest., 83:1089-1094 (1989); and Madara, Textbook of Secretory Diarrhea Eds.

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Lebenthal et al, Chapter 11, pages 125-138 (1990). barrier As between apical and basolateral compartments, they selectively regulate the passive diffusion of ions and water-soluble solutes through the paracellular pathway (Gumbiner, Am. J. Physiol., 253 (Cell Physiol. 22):C749-C758 (1987)). barrier maintains any gradient generated by the οf pathways associated with transcellular route (Diamond, Physiologist, 20:10-18 (1977)).

There is abundant evidence that ZO, once regarded as static structures, are in fact dynamic and readily adapt to a variety of developmental (Magnuson et al, *Dev. Biol.*, <u>67</u>:214-224 Revel et al, Cold Spring Harbor Symp. Quant. Biol., 15 40:443-455 (1976); and Schneeberger et al, J. Cell Sci., <u>32</u>:307-324 (1978)),physiological (Gilula et al, Dev. Biol., 50:142-168 Madara et al, J. Membr. Biol., 100:149-164 (1987); 20 Mazariegos et al, J. CellBiol., 98:1865-1877 (1984); and Sardet et al, J. Cell Biol., 80:96-117 (1979)), and pathological (Milks et al, J. Cell Biol., <u>103</u>:2729-2738 (1986); Nash et Invest., 59:531-537 (1988); and Shasby et al, Am. J. 25 Physiol., <u>255</u>(Cell Physiol., <u>24</u>):C781-C788 (1988)) circumstances. The regulatory mechanisms underlie this adaptation are still not completely understood. However, it is clear that, in the presence of Ca^{2+} , assembly of the ZO is the result of 30 cellular interactions that trigger a complex cascade of biochemical events that ultimately lead to the formation and modulation of an organized network of ZO elements, the composition of which has been only WO 00/15252 PCT/US99/18842

partially characterized (Diamond, *Physiologist*, 20:10-18 (1977)). A candidate for the transmembrane protein strands, occluding, has been identified (Furuse et al, *J. Membr. Biol.*, 87:141-150 (1985)).

5 proteins have been identified cytoplasmic submembranous plaque underlying membrane contacts, but their function remains to be established (Diamond, supra). ZO-1 and ZO-2 exist as a heterodimer (Gumbiner et al, Proc. Natl. Acad. 10 Sci., USA, 88:3460-3464 (1991)) detergent-stable complex with an uncharacterized 130 kD protein (ZO-3). Most immunoelectron microscopic studies have localized ZO-1 to precisely beneath membrane contacts (Stevenson et al, Molec. 15 Cell Biochem., 83:129-145 (1988)). Two other proteins, cingulin (Citi et al, Nature (London), 333:272-275 (1988)) and the 7H6 antigen (Zhong et al, J. Cell Biol., 120:477-483 (1993)) are localized further from the membrane and have not vet 20 been cloned. Rab 13, a small GTP binding protein has also recently been localized to the junction region (Zahraoui et al, J. Cell Biol., 124:101-115 (1994)). Other small GTP-binding proteins are known to regulate the cortical cytoskeleton, i.e., rho 25 regulates actin-membrane attachment in focal contacts (Ridley et al, Cell, 70:389-399 (1992)), and rac regulates growth factor-induced membrane ruffling (Ridley et al, Cell, 70:401-410 (1992)). Based on the analogy with the known functions of 30 plaque proteins in the better characterized cell junctions, focal contacts (Guan et al, Nature,

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J. Cell Biol., <u>123</u>:1049-1053

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(1993)), it has been hypothesized that tj-associated plaque proteins are involved in transducing signals in both directions across the cell membrane, and in regulating links to the cortical actin cytoskeleton.

To meet the many diverse physiological pathological challenges to which epithelia are subjected, the ZO must be capable of rapid and coordinated responses that require the presence of a complex regulatory system. The precise 10 characterization of the mechanisms involved in the assembly and regulation of the ZO is an area of current active investigation.

There is now a body of evidence that tj structural and functional linkages exist between the actin cytoskeleton and the tj complex of absorptive cells (Gumbiner et al, supra; Madara et al, supra; and Drenchahn et al, J. Cell Biol., 107:1037-1048 (1988)). The actin cytoskeleton is composed of a complicated meshwork of microfilaments whose precise geometry is regulated by a large cadre actin-binding proteins. An example of how the state of phosphorylation of an actin-binding protein might regulate cytoskeletal linking to the cell plasma membrane is the myristoylated alanine-rich C kinase substrate (hereinafter "MARCKS"). MARCKS is specific protein kinase C (hereinafter "PKC") substrate that is associated with the cytoplasmic face of the plasma membrane (Aderem, Elsevier Sci. Pub. (UK), pages 438-443 (1992)). In its non-phosphorylated form, MARCKS crosslinks to the Thus, it is likely that the actin membrane actin. meshwork associated with the membrane via MARCKS is relatively rigid (Hartwig et al, Nature, 356:618-622

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(1992)). Activated PKC phosphorylates MARCKS, which is released from the membrane (Rosen et al, J. Exp. (1990); and Med., <u>172</u>:1211-1215 Thelen et al, Nature, 351:320-322 (1991)). The actin linked to MARCKS is likely to be spatially separated from the membrane and be more plastic. When MARCKS dephosphorylated, it returns to the membrane where it once again crosslinks actin (Hartwig et al, supra; and Thelen et al, supra). These data suggest that the F-actin network may be rearranged by a PKC-dependent phosphorylation process that involves actin-binding proteins (MARCKS being one of them).

II. Zonula Occludens Toxin ("Zot") and Zonulin

15 Most Vibrio cholerae vaccine candidates constructed by deleting the ctxA gene encoding cholera toxin (CT) are able to elicit high antibody responses, but more than one-half of the vaccinee still develop mild diarrhea (Levine et al, Infect.

20 Immun., 56(1):161-167 (1988)). Given the magnitude of the diarrhea induced in the absence of CT, it was hypothesized that V. cholerae produce other enterotoxigenic factors, which are still present in strains deleted of the ctxA sequence (Levine et al,

25 supra). As a result, a second toxin, zonula occludens toxin (hereinafter "Zot) elaborated by V. cholerae, and which contribute to the residual diarrhea, was discovered (Fasano et al, Proc. Nat.

Acad. Sci., USA, 8:5242-5246 (1991)). The zot gene
is located immediately adjacent to the ctx genes.
The high percent concurrence of the zot gene with
the ctx genes among V. cholerae strains
(Johnson et al, J. Clin. Microb., 31/3:732-733

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(1993); and Karasawa et al, FEBS Microbiology Letters, 106:143-146 (1993)) suggests a possible synergistic role of Zot in the causation of acute dehydrating diarrhea typical of cholera. Recently, the zot gene has also been identified in other enteric pathogens (Tschape, 2nd Asian-Pacific Symposium on Typhoid fever and other Salmonellosis, 47 (Abstr.) (1994)).

It has been previously found that, when tested on rabbit ileal mucosa, Zot increases the intestinal 10 permeability by modulating the structure intercellular tight junctions (Fasano et al, supra). found has been that as a consequence modification of the pericellular pathway, 15 intestinal mucosa becomes more permeable. It also was found that Zot does not affect Na+-glucose coupled active transport, is not cytotoxic, and fails to completely abolish the transepithelial resistance (Fasano et al, supra).

20 More recently, it has been found that Zot is capable of reversibly opening tight junctions in the intestinal mucosa, and thus Zot. when co-administered with a therapeutic agent, is able to effect intestinal delivery of the therapeutic agent, 25 when employed in an oral dosage composition for intestinal drug delivery (WO 96/37196, U.S. Patent 5,827,534 and U.S. Patent 5,665,389; each of which is incorporated by reference herein in their It has also been found that Zot is entirety). 30 capable of reversibly opening tight junctions in the nasal mucosa, and thus Zot, when co-administered with a therapeutic agent, is able to enhance nasal absorption of a therapeutic agent (WO 98/30211 and

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U.S. Patent 5,908,825; which is incorporated by reference herein in its entirety).

U.S. 5,864,014 and U.S. Τn Patent Patent 5,912,323; which incorporated are by reference herein in their entirety, Zot receptors from CaCo2 cells, heart, intestinal and brain tissue has been identified and isolated. The Zot receptors represent the first step of the pericellular pathway involved in the regulation of epithelial intestinal and nasal permeability.

In U.S. Patent 5,945,510, which is incorporated reference herein in its entirety, mammalian proteins that are immunologically and functionally related Zot, and that function the to modulator physiological of mammalian tight junctions, have been identified and purified. These mammalian proteins, referred to as "zonulin", are useful for enhancing absorption of therapeutic agents across tight junctions of intestinal nasal mucosa, as well as across tight junctions of the blood brain barrier. These proteins are further characterized by the ability to bind to the receptors.

In pending U.S. Patent Application No. 09/127,815 August 25 filed 3, 1998, entitled "Peptide Antagonists of Zonulin and Methods for Use of the Same", which is incorporated by reference herein in its entirety, peptide antagonists of zonulin have been identified. Said peptide 30 antagonists bind to Zot receptor, yet do not function to physiologically modulate the opening of mammalian tight junctions. The peptide antagonists competitively inhibit the binding of Zot and zonulin WO 00/15252 PCT/US99/18842

to the Zot receptor, thereby inhibiting the ability of Zot and zonulin to physiologically modulate the opening of mammalian tight junctions.

5 III. Antigen Presenting Cells and Immune Responses

For a complete discussion of immune responses immunomodulation, see Chapter 10 "Recent and Immunology", by Sztein et al, New Advances in Generation of Vaccines, pages 99-125, Eds. (1997), the disclosure of which Levine et al

10 Levine et al (1997), the disclosure of which is hereby incorporated by reference.

One of the primary mechanisms of protection against infectious agents involves specific In contrast to innate immunity, acquired immunity. 15 the effector mechanisms of acquired immunity that include, among others, antibodies, cytotoxic "CTL"), (hereinafter lymphocytes T lymphocyte-derived cytokines (such as IFN-γ, IL-4, etc.) are induced following exposure to antigens or infectious agents and increase in magnitude with 20 successive exposures to the specific antigens. ability to "recall" previous exposures to antigens respond rapidly with immunological effector of increased magnitude (immunologic responses 25 constitutes the foundation for memory) infectious immunoprophylactic vaccination against The chief cell types involved in specific immune responses are T and B lymphocytes.

B lymphocytes or B cells are derived from the bone marrow and are the precursors of antibody secreting cells (plasma cells). B cells recognize antigens (proteins, carbohydrates or simple chemical groups) through immunoglobulin receptors on the cell

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membrane (Fearon et al, Science, 272:50-53 (1996); Ziegler-Heitbroack et al, Immunol. Today, 14:121-152 al, Adv.Immunol., Banchereau et (1993); and 52;125-262 (1992)). After triggering by antigen, they clonally expand and switch their expression of antibody isotype (e.g., IgM to IgG, IgE influence of cytokines derived from under the macrophages and other cell types. T cells, affinity В cells Somatically-mutated, high generated and selected by antigen in and around the germinal centers that are formed in lymph nodes, more disorganized and spleen, patches Peyers' the peripheral lymphoid lymphatic aggregates of system (Banchereau et al, (1996) supra; Clark et al, Immunol., 9:97-127 (1991);Ann. Rev. MacLennan et al, *Immunol*. *Today*, <u>14</u>:29-34 (1993)). They are the basis for B cell memory.

T cells, in contrast lymphocytes or to B cells, recognize peptides derived from protein antigens that are presented on the surface "APC") (hereinafter antigen presenting cells II major I or Class conjunction with Class histocompatibility complex (MHC) molecules. Clones T cell receptors lymphocytes expressing of appropriate affinity are (hereinafter "TCR") triggered by antigen to proliferate and develop into effector cells (Fearon, (1996) supra; Sprent et al al, (1994); and Hendrick Cell. <u>76</u>:315-322 ed., Germain, Fundamental Immunology, 3rd (1993)). After elimination of the pages 629-676 infectious agent, the antigen-specific clones remain as memory T cells that, upon subsequent exposures to more rapid and antigen, provide a stronger,

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sometimes qualitatively different specific immune response.

There are two main populations of T cells, those expressing CD4 molecules and those expressing CD8 molecules. CD4 and CD8 molecules are T cell surface glycoproteins that serve as important accessory molecules (co-receptors) during antigen presentation by binding to Class II and Class I MHC respectively (Hendrick molecules, et al, (1993)). Thus, CD4 and CD8 molecules significant role in stabilizing the interactions of T cells and APC initiated by the specific binding of the TCR complex to antigenic peptides presented in association with MHC molecules. Consequently, CD4 and CD8 molecules, originally used primarily as markers identify T cell to populations different functional characteristics, play a major in Class II MHC-restricted and Class MHC-restricted T cell activation. CD4+ cells (T helper or Th) are mainly involved in inflammatory responses and providing help for antibody production by B cells, while CD8+ cells (T cytotoxic or Tc) compose the majority of CTL primarily involved in Class I MHC-restricted killing of target cells infected by pathogenic including organisms, bacteria, viruses and parasites (Sztein et al, J. Immunol., <u>155</u>:3987-3993 (1995); Kaufman, Rev. Immunol., 11:129-163 (1993) and Immunol. Today, 9:168-174 (1988); Townsend et al, Cell, 44:959-968 (1986); Malik et al, Proc. Natl. Acad. Sci., USA, 88:3300-3304 (1991); Sedegah et al, J. 149:966-971 (1992); and Shearer et al, Today, 17:21-24 (1996)).

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Successful antigen specific activation T cells resulting in cell expansion and differentiation (or lymphocyte proliferation) requires a first signal provided by the interaction of TCR on the surface of T cells with MHC-antigen complexes on APC and a second, complementary, signal factors, such provided by soluble as (a co-stimulatory molecule) binding of CD28 members of the B7 family (e.g., CD80 (B7-1) or CD86 (Lenschow, Ann. (B7-2)) on APC Rev. Immunol.. 14:233-258(1996); and Linsley et al, Ann . Immunol., <u>11</u>:191-212(1993)). The study of the CD28/B7 co-stimulatory pathway and other adhesion molecules that help stabilize \mathbf{T} cell-APC interactions (and which also appear to play critical roles in lymphocyte homing), is one of the key areas in which many significant advances have been made in recent years.

Presentation of antigens to T cells involves a 20 series of intracellular events within the APC, including the generation of antigenic peptide MHC fragments, binding of these peptides molecules to form stable peptide-MHC complexes and transport of these complexes to the cell surface 25 where they can be recognized by TCR in the surface of T cells. Evidence has accumulated for the existence of two main pathways of antigen processing and presentation ("classical pathways"). One of these pathways, "cytosolic pathway", the 30 predominantly used for presentation of peptides produced endogenously in the APC, such as viral proteins, tumor antigens and self-peptides, associated with Class I MHC molecules

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(Hendrick et al, supra; and Germain, supra (1993)). The presentation of large numbers of self-peptides complexed to Class I MHC molecules results from the inability of APC to differentiate between self and Under normal conditions, most T cells non-self. 5 selected to recognize self-peptides are eliminated during T cell differentiation or are actively down regulated, and consequently can not be activated by self-peptide-Class I MHC complexes. The second antigen processing "classical pathway" of 10 presentation, "endosomal pathway", which is predominantly used for presentation of soluble exogenous antigens bound to Class II MHC molecules, involves the capture of antigen by APC, either by binding to a specific receptor or by uptake in the 15 fluid phase by macropinocytosis (Lanzavecchia, Curr. Opin. Immunol., 8:348-354 (1996)). Triggering of T cells through the TCR has been shown with as few as 200-600 peptide/MHC complexes in the case of Semin. et al, nucleoproteins (Falk 20 influenza (1993)). most immune Immunol.. 5:81-94 In associated with responses, antigenic epitopes Class I MHC molecules trigger the activation of CD8+ CTL responses, while antigenic fragments (epitopes) derived from soluble proteins complexed to Class II 25 MHC molecules are recognized by CD4+ Th cells. the most important These findings are among contributions made over the past few years on the mechanisms involved in the early stages of immune activation and are critical for the development of 30 successful vaccines.

As mentioned above, there are two "classical" pathways of antigen processing and presentation.

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The Class I MHC pathway is that most commonly used for processing of cellular proteins present in most, if not all, cellular compartments, including the cytosol, nucleus and mitochondria (Falk et al, supra (1993)) for recognition by CD8+ CTL. The Class II MHC pathway is predominantly used for processing and presentation of exogenous antigens, such as proteins produced extracellular bacteria and by infectious microorganisms that can be presented to CD4+ Th cells. Both Class I and II MHC molecules bind peptide antigens through the use of surface "receptors" or "binding clefts". However, the route antigen processing and preparation dramatically between the two. Class I antigens are processed and prepared by the "cytosolic pathway". Specifically, peptides synthesized intracellularly are degraded into small protein fragments which are then carried across the membrane of the endoplasmic reticulum (ER). Inside the ER, antigenic fragments bind to Class I MHC molecules forming a complex that is then transported to the Golgi apparatus ultimately to the cell surface where they recognized by TCR, signalling antigen-specific CTL expansion and differentiation, the first step of an immune response. Class II antigens, on the other hand, are processed and prepared by the "endosomal pathway". Specifically, native antigens captured by a circulating APC, the antigen binding to a specific or nonspecific receptor. The antigen is then internalized by the APC by a mechanism of receptor-mediated endocytosis or pinocytosis. internalized antigen is then localized endosome, a membrane bound vesicle involved in the

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degradation of intracellular transport and Cleaved peptide fragments then bind to antigen. Class II MHC molecules to form a complex that is transported through the Golgi apparatus, into the endosomal compartment, and to the cell surface to become recognized by TCR, again signaling the antigen-specific Th cell expansion and differentiation.

APC play a vital role in the generation of an 10 immune response. For presentation of processed antigens to CTL in a Class I-restricted fashion, the APC must express Class I MHC molecules and have the ability to express on the cell surface endogenously proteins complexed to Class produced Almost all cells endogenously producing 15 molecules. viral, parasitic, or bacterial proteins or tumor the cytosol antigens that gain access to function as APC. For presentation of processed Th cells in a Class II restricted antigens to 20 fashion, the APC must be able to recognize and bind antigen through specific ornonspecific receptors for the particular antigen. Cells that most efficiently present antigens to Th lymphocytes, professional APC include dendritic called cells (DC), macrophages, B lymphocytes, Langerhans 25 cells, and, in certain instances, human endothelial cells (Lanzavecchia, supra (1996)).

that originate in the bone DC marrow are considered to be the most efficient APC presentation of soluble antigens. DC capture antigens on the periphery and migrate to the spleen or lymph nodes, where they efficiently activate the Th cells, particularly naive T cells (Lanzavecchia,

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supra (1996); and Peters et al, Immunol. Several unique characteristics <u>17</u>:273-278 (1996)). enable DC to function so effectively as antigen presenters. Specifically, they have the ability to internalize soluble antigens by several mechanisms, 5 macropinocytosis, constitutive including antigen-antibody complexes internalization of through CD32 receptor binding, and internalization mannosylated or fucosylated antigens through mannose receptor binding. This allows DC to sample 10 large amounts of fluid in short periods of time, compartment lysosomal in accumulating them a containing Class II MHC molecules and proteases. a number constitutively express costimulatory and other adhesion molecules that are 15 upregulated by proinflammatory cytokines such as IL-1 α , IL-1 β , and TNF- α , thereby enhancing their ability to function as APC for Class ΙI MHC restricted Th immune responses.

Macrophages and other mononuclear phagocytes 20 are probably the most effective APC for antigens derived from most pathogenic microorganisms other than viruses through their ability to phagocytose large particles, such as bacteria and parasites. conditions, phagocytized 25 Under typical microorganisms are then killed in the phagolysosomes of generation and digested, resulting in the for available binding to fragments antigenic Class II MHC molecules for presentation to Th cells. Other important mechanisms that allow macrophages to 30 serve as effective APC include their ability to internalize soluble antigens through binding antigen-antibody complexes to CD16, CD32 and CD64

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receptors. Macrophages, also internalize complement coated proteins through receptors for C3 and other C' components and upon stimulation by growth factors, by macropinocytosis. Moreover, macrophages express receptors for mannose and are a major source of pro-inflammatory cytokines including IL-1 α , IL-1 β , IL-6, IL-8, IL-12, TNF- α , and TNF- β that exert potent immunoregulatory activities on T cell responses (Sztein et al, supra (1997)).

are very effective APC lymphocytes for 10 soluble antigens for presentation to Th cells. This is largely based on their ability to bind and internalize specific soluble antigens efficiently through the B-cell receptor complex (BCR), consisting of the specific membrane 15 immunoglobin (mIg) and the $Ig\alpha$ (CD79 α)- $Ig\beta$ (CD79 β) heterodimer (Falk et al, supra (1993)).

Langerhans cells (LC), derived from bone marrow progenitors, are considered to be the only cells present in the epidermis with APC capabilities. LC migrate out of the epidermis via the lymphatics to the regional lymph nodes where they develop into DC. Interestingly, LC express CD1, a nonclassical MHC molecule capable of presenting to T cells, in a restricted fashion, nonprotein antigens such as microbial lipid and glycolipid antigens.

The invention herein focuses on the antigen specific down-regulation of APC-mediated immune responses. The invention stems from the discovery of a macrophage surface receptor to which Zot binds in a specific and saturable way. The present invention describes a method for using Zot or zonulin as antigen-specific immunoregulators and in

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immunotherapeutics. Specifically, both Zot and zonulin inhibit APC-mediated antigen-specific lymphocyte proliferation in a dose dependent manner without affecting mitogen induced responses. This down-regulation of the immune response is at least in part associated with the decreased uptake of antigen.

Currently available of modulators immune cyclosporin and steroidal responses, such as compounds, have a generalized effect on antigen and of the immune system stimulations (Reed et al, J. Immunol., 137:150-154 (1986)). invention disclosed herein offers the advantage of enabling the down-regulation of immune responses to a particular antigen without inducing negative side susceptibility effects, such as increased immune suppression, infection and generalized typical of the immunomodulators of the prior art.

20 <u>SUMMARY OF THE INVENTION</u>

It is a object of the invention to provide a method for down-regulating an animal host's immune response to certain antigens, thereby facilitating immune based therapies. Specifically, it is an object of the invention to inhibit the ability of antigen presenting cells (APC) to process and present antigens to lymphocytes, thereby suppressing the lymphocyte proliferation and subsequent immune system reactions in response to defined antigens.

It is a further object of the invention to provide a treatment for an animal afflicted with an autoimmune or immune related disease or disorder such as multiple sclerosis, rheumatoid arthritis,

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insulin dependent diabetes mellitus, celiac disease, Sjogren's syndrome, systemic lupus erythematsosus, auto-immune thyroiditis, idiopathic thrombocytopenic hemolytic anemia, Grave's disease, Addison's disease, autoimmune orchitis, pernicious anemia, vasculitis, autoimmune coagulopathies, myasthenia gravis, polyneuritis, pemphiqus, rheumatic carditis, polymyositis, dermatomyositis, and scleroderma by administering an effective amount of a Zot-related immunoregulator. In an alternative embodiment. the treatment of the animal afflicted with an autoimmune or immune related disease disorder may involve the administration effective amount of a Zot-related immunoregulator in combination with a specific auto-immune antigen(s).

It is a further object of the invention to provide a treatment of an animal afflicted with immune rejection subsequent to tissue or organ transplantation by administering an effective amount of a Zot-related immunoregulator. In an alternative embodiment, the treatment of the animal afflicted with immune rejection subsequent to tissue or organ transplantation may involve the administration of an effective amount of a Zot-related immunoregulator in combination with а specific transplantation antigen(s).

It is a further object of the invention to provide a treatment for an animal afflicted with an inflammatory or allergic disease or disorder such as asthma, psoriasis, eczematous dermatitis, Kaposi's sarcoma, multiple sclerosis, inflammatory bowel disease, proliferative disorders of smooth muscle

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cells, and inflammatory conditions associated with mycotic, viral, parasitic, or bacterial infections by administering a therapeutically effective amount of a Zot-related immunoregulator. In an alternative embodiment, the treatment of the animal afflicted with an inflammatory or allergic disease or disorder may involve the administration of an effective amount of a Zot-related immunoregulator in combination with a specific inflammatory related antigen(s) or allergen(s).

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 illustrates Zot-FITC saturation binding curves for lymphocytes and macrophages. The data demonstrate that Zot binds preferentially to human monocytes/macrophages.

Figure 2 illustrates blocking of Zot-FITC binding by unlabeled Zot. Preincubation of PBMC with unlabeled Zot decreased the binding of Zot-FITC to both, monocyte/macrophages and T lymphocytes by about 33%, suggesting that Zot binding to these cells is receptor-mediated. Preincubation of cells with purified MBP had no effect in blocking Zot-FITC binding, indicating that blocking with unlabeled Zot is a specific phenomenon.

Figure 3 illustrates the effects of Zot on proliferation of human PBMC induced by PHA and tetanus toxoid. The data demonstrate that Zot markedly suppresses tetanus toxoid-induced proliferation in a dose dependent manner while having no effect on PHA induced proliferation

Figure 4 illustrates the effects of anti-Zot antiserum on Zot-induced suppression of

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proliferation of human PBMC induced by tetanus toxoid. Addition of anti-Zot reverses the Zot-mediated suppression of tetanus toxin-induced proliferation by greater than 50%.

5 Figures 5A-5D illustrates the effect of Zot on FITC-dextran uptake by normal human CD14+HLA-DR+ macrophages. Figure 5A depicts FITC-dextran uptake in media at 0°C (2.9%) and represents the temperature dependence of antiqen uptake. Figure 5B depicts 10 FITC-dextran uptake in media at 37°C (46.0%) and represents a control for antigen uptake. Figure 5C depicts FITC-dextran uptake in BSA at 37°C (39.0%) and represents a negative control of antigen uptake. Figure 5D depicts FITC-dextran uptake in Zot at 37°C 15 (19.3%). The data show that Zot decreases the uptake of antigen.

Figure 6 illustrates the number of FITC-Zot binding sites/cell in human macrophages lymphocytes. PBMC were incubated with increasing concentrations of Zot-FITC and analyzed by flow The mean fluorescence intensity of each cytometry. population following incubation with Zot-FITC were converted to number of Zot binding sites/cell using a standard curve constructed using the Quantum 26 MESF kit. These data show that binding of Zot is a saturable phenomenon, with saturation reached at approximately 0.5 pM and that the average number of binding sites/cell is approximately 10-fold higher in macrophages (~106,000) than in lymphocytes (~9,000).

Figures 7A-7C illustrate the kinetics of Zot binding to human macrophages and lymphocytes. Determination of the kinetics of binding of Zot to

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human cells was carried out by flow cytometry using Zot-FITC conjugates. PBMC labeled with anti-CD3-ECD and anti CD-14-PE mAb were maintained at 37°C for the duration of the experiment (12 min) while data was collected using a viable sample handler (kinetics module) attached to the flow cytometer. FITC fluorescence levels were collected 90-150 seconds (indicated by the arrows) and data acquisition was paused for ~10-15 sec to inject Zot-FITC (Figures 7A and 7B). Figure 7C shows the kinetics of binding of anti-CD14-FITC to unlabeled human monocyte/macrophages. Data are presented as isometric displays of Zot-FITC intensity (y axis) versus time (x axis) versus cell number (z axis) for gated onCD3 (lymphocytes) or CD14 (macrophages). The results indicate that Zot binding to human macrophages (Figure 7A) and lymphocytes (Figure 7B) occurs very rapidly, reaching equilibrium within 2 min following addition of Zot-FITC.

Figure 8 illustrates the binding of FITC-Zot to human T and B lymphocytes. PBMC were incubated with Zot-FITC and mAb to molecules present in T (CD3 $^+$) and B (CD19 $^+$) lymphocytes and analyzed by flow cytometry.

25 An isotopic FITC-labeled control mAb (mIg) corresponding to cells gated on the forward vs. side scatter lymphocyte region is also shown as an indicator of non-specific binding. The results indicate that Zot binds to both T and B lymphocytes.

Figure 9 illustrates the inability of Zot antagonists to block Zot-FITC binding. PBMC stained with CD14-PE and CD3-ECD were washed and incubated for 15 min at 4°C in AIM-V medium alone or with the

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addition of 100-fold excess of FZI/0 (SEQ ID NO:7), FZI/1 (SEQ ID NO:8), BSA (negative control) or 4-fold excess of unlabeled Zot (positive control). Cells were then incubated with Zot-FITC and analyzed by flow cytometry. The results are expressed as % suppression of the mean fluorescence intensity of cells incubated with Zot-FITC in the presence of Zot antagonists, unlabeled Zot or BSA as related to the mean fluorescence intensity of cells incubated in media alone (arbritarily assigned a value of 100%). The results show that the addition of either FZI/0 or FZI/1 Zot antagonists did not significantly block binding of Zot-FITC to CD14⁺ gated macrophages. contrast, pre-incubation with unlabeled Zot blocked binding of Zot-FITC by 24-43%.

Figures 10A-10B illustrate Zot suppression of CD14 expression on human monocytes/macrophages. PBMC were incubated for 18 hr in the absence or presence of TT without or with purified Zot or BSA, stained with CD14-FITC and analyzed by cytometry. The results are shown as single color histograms of CD14 fluorescence on cells gated on the "monocyte region", defined based on the forward scatter vs. side scatter characteristics of human macrophages. The addition of Zot caused a marked suppression of the expression of CD14 in the absence of TT (Figure 10A) or presence of TT (Figure 10B).

Figure 11 illustrates the effects of Zot on human monocytes/macrophages and lymphocyte viability. PBMC were incubated for various time periods in the absence or presence of purified Zot or BSA. Cell viability was assessed using the propidium iodide exclusion test and flow cytometry.

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The results are shown as the % viable cells gated on the "monocyte region" or "lymphocyte region", defined based on the forward scatter vs. side scatter characteristics of these cell populations. The results show that the addition of Zot affect macrophage viability at relatively early times, while the effects on lymphocytes did not become apparent until at least 4 days in culture.

Figures 12A-12C illustrates Zot-mediated 10 induction of cytokine production by human monocytes/macrophages. PBMC were incubated for 6 hr to 4 days in the absence or presence of purified Zot or BSA. Supernatants were collected indicated times and cytokine levels measured by 15 chemiluminescence ELISA. Addition of Zot resulted the production of high levels of $TNF-\alpha$ (Figure 12A) and IL-10 (Figure 12C) as early as 6 hr, reaching peak levels at 24 hr. A weak induction of IL-1 β production was also observed 20 (Figure 12B).

13A-13B Figures illustrate Zot-mediated induction of cytokine production by human lymphocytes. PBMC were incubated for 3 days without or with TT in the absence or presence of purified Zot or BSA and cytokine levels in the supernatants measured by chemiluminescence ELISA. Addition of Zot resulted in the suppression of IL-2 production induced by incubation with TT, while no measurable levels of IL-2 were induced by Zot in the absence of TT (Figure 13A). In contrast, addition of Zot consistently induced the production of IFN-y in the absence of TT, similar to the levels induced by TT,

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and markedly increased the levels of IFN- γ induced by TT (Figure 13B).

DETAILED DESCRIPTION OF THE INVENTION

Previous studies have focused on the ability of and zonulin to physiologically modulate the opening of mammalian tight junctions (or zonula occludens) of the epithelia of various tissues, such modulation particularly useful to facilitate drug delivery across these membranes. In the course of study, receptors to Zot and zonulin were identified and isolated from CaCo2 cells, heart, intestinal and brain tissue. Further to discovery of Zot receptors, peptide antagonists of zonulin were identified, said peptide antagonists binding to Zot receptor, yet not functioning to physiologically modulate the opening of mammalian tight junctions (i.e., lacking the biologic activity). The peptide antagonists competitively inhibit the binding of Zot and zonulin to the Zot receptor, thereby inhibiting the ability of Zot and zonulin to regulate the tight junctions.

In light of the known effect of Zot and zonulin on the paracellular pathway, it was surprising to find a receptor for Zot on fully differentiated macrophages isolated from blood. was initially unclear why circulating cells such as macrophages would have a receptor for a molecule associated with tissue cell modulation. In pursuing the answer to this question, it was discovered that Zot and zonulin also have the ability to physiologically regulate the activity macrophages. Though not wishing to be bound by

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theory, it appears that Zot is blocking the receptor on the macrophage, binding to the macrophage surface receptor in a specific and saturable way. binding to macrophages, Zot alters the macrophages and present ability to process an antigen lymphocytes, ultimately suppressing the proliferation of lymphocytes in response to the antigen in a dose dependent and antigen-specific manner. In other words, Zot allows antigen-specific down-regulation of an immune The results described in detail below response. further suggest that this down-regulation of the immune response is at least in part associated with the decreased uptake of antigen. It appears that Zot is a multi-functional protein, controlling the immune response by a dual mechanism of regulating the uptake and the trafficking of antigens.

Zonula occludens toxin or "Zot" is produced by The particular strain of V. cholera V. cholerae. 20 from which Zot is derived is not critical to the Examples of such V. cholerae present invention. strains include 569B. 395 E7946 strain and (Levine et al, supra; Johnson et al, supra; and Karasawa et al, supra).

25 As used herein, "Zot" refers to the mature protein of 399 amino acids, as well as mutants thereof which retain the ability to regulate tj. For example, an N-terminal deletion of amino acids 1-8 can be made without effecting 30 activity, and N-terminal fusion proteins of Zot can made without effecting Zot activity. mutants can be readily prepared by site-directed

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mutagenesis, and screened for Zot activity as described herein.

Zot can be obtained and purified, e.g., genetically-engineered E. coli strains over-expressing the zot gene (Baudry et al, Infect. Immun., 60:428-434 (1992)), alone or fused to other genes, such as maltose binding protein below), glutathione-S-transferase Example 1 (see Example 2 below), or 6 poly-histidine (see Example 2 below).

As used herein, the term "zonulin" refers to a substantially pure biologically active protein having an apparent molecular weight of about 47 kDa, determined by SDS-polyacrylamide 15 electrophoresis, and the following N-terminal amino acid sequence: Asn Asp Gln Pro Pro Pro Ala Gly Val Thr Ala Tyr Asp Tyr Leu Val Ile Gln (SEQ ID NO:1), or the following N-terminal amino acid sequence: Glu Val Gln Leu Val Glu Ser Gly Gly Xaa Leu Val Gln 20 Pro Gly Gly Ser Leu Arg Leu (SEQ ID NO:2) as well as mutants thereof which retain the ability bind to the receptor for Zot and to regulate tj.

Zonulin is produced by, or found in, various mammalian cells and tissues, e.g., rabbit or human cells/tissue. The particular mammalian cell/tissue type from which zonulin is derived is not critical to the present invention. Examples of such mammalian tissue types include heart, lung, intestine, liver, brain, kidney, and pancreas.

Zonulin can be obtained and purified, e.g., by affinity-purification chromatography using anti-ZOT antibodies, as described in Example 3 of U.S. Patent 5,945,510, incorporated by reference herein.

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terms "Zot-related herein, the used immunoregulator" or "Zot-related immunoregulating molecule" refers to Zot or zonulin, as defined as used herein, the term Furthermore, above. "inhibition" (and "inhibit" and "inhibiting") refers to the down regulation of any aspect of APC activity substantial reduction results in а proliferation. Example activities lymphocyte include antigen uptake, antigen processing, antigen presentation. The term "inhibition" as used necessarily imply complete herein does not suppression of function or result. Rather, partial inhibition is contemplated by the term.

To provide a treatment for an animal afflicted with an autoimmune or immune related 15 disease or disorder such as multiple sclerosis, insulin dependent diabetes rheumatoid arthritis, celiac disease, Sjogren's syndrome, mellitus, erythematsosus, auto-immune lupus systemic idiopathic thrombocytopenic purpura, 20 thyroiditis, hemolytic anemia, Grave's disease, Addison disease, autoimmune orchitis, pernicious anemia, vasculitis, myasthenia coagulopathies, gravis, autoimmune rheumatic carditis, polyneuritis, pemphiqus, 25 polymyositis, dermatomyositis, and scleroderma, a Zot-related immunoregulator is administered alone or in combination with a specific auto-immune related Examples of specific auto-immune antigens antigen. associated with auto-immune diseases or disorders are gliadin (antigen associated with celiac disease) 30 and myelin basic protein (associated with multiple sclerosis).

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To provide a treatment for an animal host afflicted with immune rejection subsequent to tissue or organ transplantation, a Zot-related immunoregulator is administered alone or in combination with a specific transplantation antigen. Transplantation antigens may be obtained by assaying for the HLA of the transplant tissue or organ.

To provide a treatment for an animal afflicted with an inflammatory or allergic disease or disorder such as asthma, psoriasis, eczematous 10 dermatitis, Kaposi's sarcoma, multiple sclerosis, inflammatory bowel disease, proliferative disorders of smooth muscle cells, and inflammatory conditions mycotic, viral, associated with parasitic, bacterial infections, a Zot-related immunoregulator 15 administered alone or in combination with a specific inflammatory related antigen or allergen. Examples of specific inflammatory related antigens associated with inflammatory or allergic diseases or 20 disorders are pollens and dust (associated with asthma), proteins found in cow's milk or fragments (associated with eczematous dermatitis), thereof protein (associated with multiple myelin base and vaccine antigens to the particular sclerosis) 25 virus, parasite or bacteria associated with the inflammatory condition.

The present invention allows for the antigen-specific down-regulation of the immune response. As discussed above, one embodiment of the present invention involves the administration of an effective amount of a Zot-related immunoregulator (Zot or zonulin) alone or in combination with a specific antigen, the lymphocyte response to said

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specifically suppressed antigen being in dose-dependent manner. It is clear from the results the details of which are presented herein that the present invention is limited to those antigens processed and presented by antigen presenting cells (APC) those requiring macrophage mediation. invention relates to all APCs including macrophages and other mononuclear phagocytes, dendritic cells, Langerhans cells and B lymphocytes, endothelial cells. In a preferred embodiment, the APC is a macrophage.

The present invention may be utilized both in vivo or in vitro environments. The only criticality is the administration to animal cells, said cells either in a living host or in a cell culture. Animal cells are defined as nucleated, non-chloroplast containing cells derived from or present in multicellular organisms whose taxonomic position lies within the kingdom animalia, a primary cell culture, explant culture or a transformed cell line.

The recipient or host animals employed in the present invention are not critical thereto and include cells present in or derived from all organisms within the kingdom animalia. In preferred embodiment, the is within the animal family of mammals. Preferred animal and animal cells are mammal cells, such as humans, bovine, ovine, porcine, feline, buffalo, canine, equine, donkey, deer and primates. The most preferred animal or animal cells are human or human cells.

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ofand method particular mode The administration is not critical to the invention. The only criticality is that both the Zot-related immunoregulating molecule and the antigen reach the macrophage intact. In the context of the present 5 invention, the Zot-related immunoregulating molecule with the antigen. co-administered may be two may be administered Alternatively, the sequentially. For simplicity sake, discussion of of administration and pharmaceutical 10 preparation below are directed to administration and preparation of the antigen. However, it is clear that the same modes apply to the administration of Zot-related immunoregulating molecule. the 15 Furthermore, though the discussion is limited to administering a single antigen, it is clear that more than one antigen can be administered, immune response to more than one antigen being subsequently down-regulated simultaneously.

Successful administration requires the delivery 20 Preferred rich environment. to an APC administration routes include those that sites of the immune system, such as the mucosa or Thus, intranasal, intraocular, the lymph tissues. intraintestinal, and intravaginal are preferred 25 administration routes. This does not preclude that intradermal, parenteral administration, such as intramuscular, subcutaneous and intravenous, might also be effective routes of administration of Zot or zonulin, alone or in combination with the preferred 30 antigen(s).

Depending on the particular administration route, the dosage form may be solid, semisolid, or

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liquid preparation. The dosage form may include those additives, lubricants, stabilizers, buffers, coatings, and excipients as is standard in the art of pharmaceutical formulations.

mode of administration, the 5 Regarding the be administered oral as dosage antigen can compositions for small intestinal delivery. dosage compositions for small intestinal oral delivery are well-known in the art, and generally tablets comprise gastroresistent orcapsules 10 (Remington's Pharmaceutical Sciences, 16th Ed., Eds. Mack Publishing Co., Chapter 89 (1980);Digenis et al, *J. Pharm. Sci.*, <u>83</u>:915-921 (1994);Vantini et al, Clinica Terapeutica, 145:445-451 (1993); Yoshitomi et al, Chem. Pharm. Bull., 15 <u>40</u>:1902-1905 (1992);Thoma et al, Pharmazie, 46:331-336 (1991); Morishita et al, Drug Design and Delivery, <u>7</u>:309-319 (1991); and Lin et al, Pharmaceutical Res., 8:919-924 (1991)); each of which is incorporated by reference herein in its 20 entirety).

Tablets are made gastroresistent by the addition of, e.g., either cellulose acetate phthalate or cellulose acetate terephthalate.

Capsules are solid dosage forms in which the antigen is enclosed in either a hard or soft, soluble container or shell of gelatin. The gelatin used in the manufacture of capsules is obtained from collagenous material by hydrolysis. There are two types of gelatin. Type A, derived from pork skins by acid processing, and Type B, obtained from bones and animal skins by alkaline processing. The use of hard gelatin capsules permit a choice in prescribing

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a single antigen or a combination thereof at the considered best for dosage level individual subject. The hard gelatin consists of two sections, one slipping over other, thus completely surrounding the alone or in combination with the immunoregulating These capsules are filled by introducing molecule. the antigen, or gastroresistent beads containing the antigen, into the longer end of the capsule, and then slipping on the cap. Hard gelatin capsules are made largely from gelatin, FD&C colorants, sometimes an opacifying agent, such as titanium The USP permits the gelatin for this dioxide. purpose to contain 0.15% (w/v) sulfur dioxide to prevent decomposition during manufacture.

In the context of the present invention, oral dosage compositions for small intestinal delivery also include liquid compositions which contain aqueous buffering agents that prevent the antigen from being significantly inactivated by gastric fluids in the stomach, thereby allowing the antigen to reach the small intestines in an active form. Examples of such aqueous buffering agents which can be employed in the present invention include bicarbonate buffer (pH 5.5 to 8.7, preferably about pH 7.4).

When the oral dosage composition is a liquid composition, it is preferable that the composition be prepared just prior to administration so as to minimize stability problems. In this case, the liquid composition can be prepared by dissolving lyophilized peptide antagonist in the aqueous buffering agent.

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Likewise, other dosage delivery vehicles are contemplated by the present invention including but not limited to liposomes, cochleates, water soluble polymers and microspheres. The dosage composition may further include adjuvants such as monophosphoryl lipid A, QS-21, ISCOMs, and cytokines.

administered also be can antigen intravenous dosage compositions for delivery to the immune system. Such systemic elements of the in art. the and compositions are well-known physiological generally comprise a compositions diluent, e.g., distilled water, or 0.9% (w/v) NaCl. Likewise, the administration may be parenteral, intramuscular, or subcutaneous and intradermal, for such mentioned Dosage forms above. administration would clearly include including pharmaceutically form, acceptable physiologic buffers, diluents or the like.

an effective amount of As used herein, Zot-related immunoregulator, such as Zot or zonulin, 20 refers to an amount effective to down-regulate the activity of said antigen presenting cell, thereby antigen down-regulate the effective to presenting cell-mediated lymphocyte proliferation. Zot-related specific amount of antigen and 25 immunoregulator molecule employed is not critical to the present invention and will vary depending upon the disease or condition being treated, as well as age, weight and sex of the subject being the such a final achieve Generally, to 30 treated. concentration in, e.g., the intestines or blood, the amount of Zot-related immunoregulator molecule in a dosage composition of the present single oral

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invention will generally be about 0.1 μg to about 100 μ g, preferably about 2.0 μ g to about 60 μ g, more preferably about 20 µg to about 50 µg. the amount of antigen in a single oral dosage composition of the present invention will generally be in the range of about 0.01 μg to about 1000 μg , more preferably about 0.1 µg to about Obviously, the exact dosage of antigen will vary with the disease or disorder being treated, the preferred ranges being readily determinable through optimization experimentation routine and proceedings.

provided for following examples are The purposes only, and are in illustrative of the limit the scope 15 intended to invention.

EXAMPLE 1 Binding of FITC-labeled Zot to Lymphocytes and Macrophages

A. Materials and Methods

Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

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by density gradient isolated PBMC were lymphocyte separation centrifugation over from healthy Organon-Teknika, Durham, NC) Donors were adults and gave informed volunteers. PBMC used were consent for the blood drawing. RPMI frozen in orwere aliquoted and fresh containing 10% (v/v) FCS and 10% (v/v) DMSO using a controlled linear rate freezer apparatus (1°C per min, Planner Biomed, Salisbury, England) to preserve cell viability and maximize cell recovery. Cells

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were stored in liquid nitrogen until used. In some experiments, cells were used immediately after isolation.

5 Preparation of Purified ZOT-MBP

5000 ml of the supernatant fraction obtained cholerae culturing V. strain (Michalski et al, Infect. Immun., G1:4462-4468 (1993), which had been transformed with plasmid pZ14, was concentrated 1000-fold using a lamina flow filter with a MW cutoff of 10 kDa. The construction of pZ14, which contains the Vibrio cholera zot gene, is described in detail in, inter alia, WO 96/37196. resulting supernatant was then subjected to 8.0% (w/v) SDS-PAGE. Protein bands were detected by Coomassie blue staining of the SDS-PAGE gel. protein band corresponding to Zot was detectable when compared to control supernatant from strain CVD110 transformed with plasmid pTTQ181 (Amersham, Arlington Heights, IL), and treated in the same Therefore, even though the zot gene was manner. placed behind the highly inducible and strong tac promoter in pZ14, the level of the protein in 1000-fold concentrated pZ14 supernatant was still not detectable by the Coomassie stained SDS-PAGE gel.

Hence, to increase the amount of Zot produced, the zot gene was fused in frame with the maltose binding protein (hereinafter "MBP") gene to create a MBP-ZOT fusion protein.

The MBP vector pMAL-c2 (Biolab) was used to express and purify Zot by fusing the zot gene to the malE gene of E. coli. This construct uses the

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tac promoter, and the malEinducible strong, translation initiation signals to give high level expression of the cloned zot gene. The vector pMAL-c2 has an exact deletion of the malE signal sequence, which leads to cytoplasmic expression of Affinity chromatography fusion protein. the facilitate used purification for MBP was to isolation of the fusion protein (Biolab).

pMAL-c2 specifically, vector More linearized with EcoRI (that cuts at the 3' end of 10 the malE gene), filled in with Klenow fragment, and digested with XbaI (that has a single site The orf encoding ZOT was pMAL-c2 polylinker). subcloned from plasmid pBB241 (Baudry et al, Infect. Immun., 60:428-434 (1992)). Plasmid pBB241 was 15 Klenow BssHII, filled in with with digested digested with XbaI. Then, the fragment, and blunt-XbaI fragment was subcloned into pMAL-c2 to give plasmid pLC10-c. Since both the insert, and the vector had blunt and sticky ends, the correct 20 orientation was obtained with the 3' end of malE pLC10-c. fused with the 5' terminus of the insert. was then electroporated into E. coli strain DH5a. In pBB241, the BssHII restriction site is within the zot orf. Thus, amino acids 1-8 of ZOT are missing 25 in the MBP-ZOT fusion protein.

In order to purify the MBP-Zot fusion protein, 10 ml of Luria Bertani broth containing 0.2% (w/v) glucose and 100 μ g/ml ampicillin were inoculated with a single colony containing pLC10-c, and incubated overnight at 37°C with shaking. The culture was diluted 1:100 in 1.0 ml of the same fresh medium, and grown at 37°C while shaking, to

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about 1.0 x 10^8 cells/ml. 0.2 mM IPTG was then added to induce the MBP-Zot expression, and the culture was incubated at 37°C for additional 3 hr. bacteria were then pelleted and resuspended in 20 ml comprising ice cold "column buffer" Tris-HCl, 0.2 M NaCl, 1.0 mM EDTA, 10 mM 2-ME, The bacterial suspension was lysed by $1.0 \text{ mM NaN}_3.$ french press treatment and spun for 30 min at The supernatant was collected, 13,000 x g at 4° C. diluted 1:5 with column buffer and loaded into a 1 X 10 column of amylose resin (Biolabs, MBP-fusion purification system), pre-equilibrated with column buffer. After washing the column with 5 volumes of column buffer, the MBP-ZOT fusion protein was eluted by loading 10 ml of 10 mM maltose in column buffer. 15 The typical yield from 1.0 ml of culture was 2-3 mg of protein.

The MBP fusion partner of the purified MBP-Zot fusion protein was then cleaved off using 1.0 μg of Factor Xa protease (Biolabs) per 20 μg of MBP-Zot. Factor Xa protease cleaves just before the amino terminus of Zot. The Zot protein so obtained was run on a 8.0% (w/v) SDS-PAGE gel, and electroeluted from the gel using an electroseparation chamber (Schleicher & Schuell, Keene, NH).

When tested in Ussing chambers, the resulting purified Zot induced a dose-dependent decrease of Rt, with an ED₅₀ of 7.5 x 10^{-8} M.

Conjugation of ZOT-MBP to fluorescein 30 isothiocyanate (ZOT-FITC)

Conjugation of Zot-MBP to FITC was performed following standard techniques. Briefly, Zot-MBP was

buffer 500 ml FITC labeling dialyzed against comprising 0.1 M bicarbonate buffer (e.g., 0.09 M NaHCO₃ + 0.0085 M Na₂CO₃), adjusted to pH 9.0 with concentrated NaOH and stored at 4°C, at 4°C for 8 hr to raise the pH to 9.0. Ten μl of 5.0 mg/ml FITC in DMSO for each milligram of MBP-ZOT was then added, followed by an overnight incubation in PBS at 4°C. Unbound FITC was then removed by dialysis in 500 ml dialysis buffer comprising PBS (pH 7.4), stored at 4°C, at 4°C with two to three changes over 2 days. This preparation was stored at 4°C until used.

Binding of FITC-ZOT to Human PBMC and Flow Cytometric Analysis

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PBMC isolated as described above were incubated Zot-FITC increasing concentrations of 60 min at 37°C in siliconized tubes (to preclude the binding of macrophages to the test tube walls) in CD14 of monoclonal antibodies to 20 presence CD3 phycoerythrin (PE) and to conjugated to dye, coupled ECD (energy conjugated to PE-Texas-Red conjugate). CD14 is a marker of human monocytes/macrophages, while CD3 is a marker of fluorochromes use of these The T lymphocytes. 25 (e.g., FITC, PE and ECD) allowed the simultaneous study of the binding of ZOT to monocytes/macrophages and T lymphocytes in mixed PBMC populations by 3-color flow cytometry. Cells were then washed twice with PBS (pH 7.2) containing 1.0% (w/v) BSA 30 and 0.1% (w/v) NaAzide, and analyzed immediately by flow cytometry using an Epics Elite flow cytometer/cell sorter system.

In these experiments, fluorochrome-labeled mAbs of the same isotypes, but irrelevant specificity, Platelets, erythrocytes were used as a controls. (if any) and cell debris were excluded from analysis by setting an appropriate gate on the forward vs. Data was collected 90% light scatter parameters. 10,000 cells. for each sample for over Epics the Elite analysis was performed using analysis package (Coulter) or the WinList list-mode analysis package (Verity Software House, Topsham, ME).

B. Results

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A representative experiment showing the flow cytometric analysis of Zot-FITC binding to human 15 T lymphocytes (CD3⁺) and monocyte/macrophages (CD14⁺) is shown in Figure 1. The results show that binding as evidenced by mean fluorescent Zot-FITC, several fold higher intensity levels, is lymphocytes. T than in 20 monocytes/macrophages amounts addition of increasing Furthermore, Zot-FITC results in increased binding that start to level off after addition of 40-60 μ l of Zot-FITC. These results indicate that Zot binds preferentially 25 to human monocyte/macrophages.

Next, the binding of Zot-FITC to human monocyte/macrophages and lymphocytes in the presence of unlabeled Zot was tested, to determine if the unlabeled Zot could block binding. As shown in Figure 2, preincubation of PBMC for 30 min at 37° C with 100 μ l of unlabeled Zot, followed by the addition of ZOT-FITC (10 μ l) for 30 min at 37° C, decreased by ~33% the binding of Zot-FITC to both,

monocyte/macrophages and T lymphocytes, suggesting cells is to these binding Zot Preincubation of cells with receptor-mediated. 100 μ l of purified MBP, followed by the addition of ZOT-FITC (10 μ l) for 30 min at 37°C, had no effect binding, indicating Zot-FITC blocking Zot is a specific unlabeled with blocking phenomenon.

10 EXAMPLE 2
Proliferative Responses to Mitogens
and Antigens By Human Mononuclear Cells

A. Materials and Methods

15 Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

gradient density by PBMC isolated were separation media lymphocyte centrifugation over Organon-Teknika, Durham, from healthy NC) 20 In accordance with the institutional volunteers. review board of University of Maryland, Baltimore, donors were adults and gave informed consent for the PBMC used were fresh or were blood drawing. aliquoted and frozen in RPMI containing 10% (v/v)25 FCS and 10% (v/v) DMSO using a controlled linear rate freezer apparatus (1°C per min, Planner Biomed, Salisbury, England) to preserve cell viability and maximize cell recovery. Cells were stored in liquid nitrogen until used. In some experiments cells were used immediately after isolation.

Preparation of purified Zot

The zot gene was amplified by PCR with Deep Vent polymerase (New England Biolabs), using pBB241

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plasmid (Baudry et al, supra) DNA as a template. The forward and reverse primers used were: 5'-CGGGATCCCGTATGAGTATCTTT-3' (SEQ IDNO:3); and 5'-CCCAAGCTTGGGTCAAAATATACT-3' ID (SEQ NO:4), respectively. The 5 ' tails of these BamHIHindIII oligonucleotides contain and а а respectively. The restriction site, resulting amplicon (1.2 kb) was analyzed by 8.0% (w/v) agarose gel electrophoresis, and purified from salts and free nucleotides using an Xtreme spin column (Pierce). The above-noted two restriction enzymes were then used to digest the purified amplicon, and the resulting digested-amplicon was then inserted in the vector pQE30 (Quiagen), which had previously digested with BamHI and HindIII, so as to obtain plasmid pSU113. pQE30 is an expression vector that provides high level expression of a recombinant protein with a 6 poly-histidine tag (6xHis). The expression product of plasmid pSU113 is therefore a 6xHis-Zot fusion protein. pSU113 was then transformed into E. coli DH5a.

order to purify the 6xHis-Zot protein, the resulting transformed E. coli were grown overnight at 37°C in 150 ml of Luria Bertani broth containing 2.0% (w/v) glucose, 25 μ g/ml of kanamycin and 200 $\mu g/ml$ of ampicillin until the A_{600} about 1.10. Next, 75 ml of the overnight was cultures were added to 1000 ml of Luria Bertani broth containing 2.0% (w/v) glucose, 25 µg/ml of kanamycin and 200 μg/ml of ampicillin, incubated for about 3 hr at 37°C, with vigorous shaking, until the A_{600} was about 0.7-0.9. Then, IPTG was added to a final concentration of 2.0 mM, and .growth was

allowed to continue for 5 hrs at 37°C. Next, the cells were harvested by centrifugation at 4000 x g for 20 min, the cells resuspend in 5.0 ml/g wet weight of buffer A comprising 6.0 M GuHCl, 0.1 M sodium phosphate, and 0.01 M Tris-HCl (pH 8.0), and stirred for 1 hr at room temperature. Then, mixture was centrifuged at 10,000 x g for 30 min at 4°C, and to the resulting supernatant was added 4.0-5.0 ml/g wet weight of a 50% slurry of SUPERFLOW resin (QIAGEN), and stirring was carried out for 10 1 hr at room temperature. The resulting resin was loaded into a 1.6 x 8.0 column, which was then sequentially with buffer Α, buffer washed comprising 8.0 M urea, 0.1 M sodium phosphate, and 0.01 M Tris-HCl (pH 8.0) and buffer C comprising 15 8.0 M urea, 0.1 M sodium phosphate, and 0.01 M Tris-HCl (pH 6.3). Each wash was carried out until the A_{600} of the flow-through was less than 0.01. 6xHis-ZOT fusion protein was eluted from the column using 20 ml of buffer C containing 250 mM imidazole. 20 Then, the fractions containing with the 6xHis-ZOT fusion protein were checked by SDS-PAGE using the procedure described by Davis, Ann. N.Y. Acad. Sci., 121:404 (1964), and the gel stained with Comassie The fractions containing 6xHis-ZOT fusion 25 blue. protein were dialyzed against 8.0 M urea, combined, and then diluted 100 times in PBS. Next, 4.0 ml of a 50% slurry of SUPERFLOW resin was added, stirring was carried out for 2 hrs at room temperature, and the resulting resin loaded into a 1.6 x 8.0 column, 30 which was then washed with 50 ml of PBS. 6xHis-Zot fusion protein was eluted from the column with 10 ml of PBS containing 250 mM imidazole.

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resulting eluant was dialyzed against PBS, and the 6xHis-ZOT fusion protein was checked by SDS-PAGE, as described above.

5 Preparation and Purification of Rabbit Anti-Zot Antiserum

To obtain specific antiserum, a chimeric glutathione S-transferase (GST)-Zot protein was expressed and purified.

More specifically, oligonucleotide primers were used to amplify the zot orf by polymerase chain reaction (PCR) using plasmid pBB241 (Baudry et al, template DNA. The forward primer supra) as (TCATCACGGC GCGCCAGG, SEQ ID NO:5) corresponded to nucleotides 15-32 of zot orf, and the reverse primer (GGAGGTCTAG AATCTGCCCG AT, SEQ ID NO:6) corresponded to the 5' end of ctxA orf. Therefore, acids 1-5 of ZOT were missing in the resulting fusion protein. The amplification product was inserted into the polylinker (SmaI site) located at the end of the GST gene in pGEX-2T (Pharmacia, Milwaukee, WI). pGEX-2T is a fusion-protein expression vector that expresses a cloned gene as a fusion protein with GST of Schistosoma japonicum. The fusion gene is under the control of the tac promoter. Upon induction with IPTG, derepression occurs and GST fusion protein is expressed.

The resulting recombinant plasmid, named pLC11, was electroporated in $E.\ coli$ DH5 α . In order to purify GST-Zot fusion protein, 10 ml of Luria Bertani broth containing 100 $\mu g/ml$ ampicillin were inoculated with a single colony containing pLC11, and incubated overnight at 37°C with shaking. The

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culture was diluted 1:100 in 1.0 ml of the same fresh medium and grown at 37°C while shaking, to about 1.0×10^8 cells/ml. 0.2 mM IPTG was then added to induce the GST-Zot expression, and the culture was incubated at 37°C for additional 3 hr. bacteria were then pelleted, resuspended in 20 ml of ice cold PBS (pH 7.4), and lysed by the french press The GST- Zot fusion protein was not soluble under these conditions as it sedimented with the bacterial pellet fraction. Therefore, the pellet was resuspended in Laemli lysis buffer comprising 0.00625 M Tris-HCl (pH 6.8), 0.2 M 2-ME, 2.0% (w/v) SDS, 0.025% (w/v) bromophenol blue and 10% (v/v) glycerol, and subjected to electrophoresis on a 8.0% (w/v) PAGE-SDS gel, and stained with Coomassie A band of about 70 kDa (26 kDa of brilliant blue. GST + 44 kDA of Zot), corresponding to the fusion protein, was electroeluted from the gel using an electroseparation chamber (Schleicher & Schuell, Keene, NH).

 $10~\mu g$ of the resulting eluted protein $(10\text{--}20~\mu g)$ was injected into a rabbit mixed with an equal volume of Freund's complete adjuvant. Two booster doses were administered with Freund's incomplete adjuvant four and eight weeks later. One month later the rabbit was bled.

To determine the production of specific antibodies, 10^{-10} M of Zot, along with the two fusion proteins MBP-Zot and GST-Zot, was transferred onto a nylon membrane and incubated with a 1:5000 dilution of the rabbit antiserum overnight at 4°C with moderate shaking. The filter was then washed 15 min 4 times with PBS containing 0.05% (v/v) Tween 20

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(hereinafter "PBS-T"), and incubated with a 1:30,000 dilution of goat anti-rabbit IgG conjugated to horseradish peroxidase for 2 hr at room temperature. The filter was washed again for 15 min 4 times with PBS containing 0.1% (v/v) Tween, and immunoreactive bands were detected using enhanced chemiluminescence (Amersham).

On immunoblot, the rabbit antiserum was found to recognize Zot, as well as MBP-Zot and GST-Zot fusion proteins, but not the MBP negative control.

Moreover, to confirm the production of appropriate anti-Zot antibodies, neutralization experiments were conducted in Ussing chambers. When pre-incubated with pZ14 supernatant at 37°C for 60 min, the Zot-specific antiserum (1:100 dilution), was able to completely neutralize the decrease in Rt induced by Zot on rabbit ileum mounted in Ussing chambers.

Next, the anti-Zot antibodies were 20 affinity-purified using an MBP-Zot affinity column. More specifically, a MBP-Zot affinity column was prepared by immobilizing, overnight at temperature, 1.0 mg of purified MBP-Zot, obtained as described in Example 1 above, to a pre-activated gel 25 (Aminolink, Pierce). The column was washed with PBS, and then loaded with 2.0 ml of anti-ZOT rabbit After a 90 min incubation at room antiserum. temperature, the column was washed with 14 ml of and the specific anti-Zot antibodies were 30 eluted from the column with 4.0 ml of a solution comprising 50 mM glycine (pH 2.5), 150 mM NaCl, and 0.1% (v/v) Triton X-100. The pH of the 1.0 ml

eluted fractions was immediately neutralized with 1.0 N NaOH.

Culture conditions and 5 <u>lymphoproliferation assays</u>

 $(1.5 \times 10^6 \text{ cells/ml})$ were cultured in 1.0 ml of complete medium, (cRPMI) comprising RPMI 1640 containing 10% (v/v) fetal calf serum and 10 50 μg/ml gentamicin. Cells were incubated at 37°C, 5% CO2 in 96-well plates in the absence or presence of phytohemagglutinin (PHA, a nonspecific mitogen, used at 2.0 $\mu g/ml$) or tetanus toxoid (TT; a specific antigen, used at 2.0 µg/ml; Connaught, Swift Water, 15 PA) without or with purified ZOT (used at 20 or 60 μg/ml) or bovine serum albumin (BSA; a control protein, used at 20 or 60 µg/ml; Fraction V, Sigma, St. Louis, MO). In some experiments, an anti-ZOT rabbit antiserum or normal rabbit serum was also 20 added to the cultures at initiation. Cells were cultured for 2 days (for PHA) or 6 days (for TT) and 1.0 μ Ci/ well of tritiated thymidine was added. Plates were harvested 20 hr later on a Wallac cell (Gaithersburg, harvester MD) and incorporated 25 thymidine measured on a Wallac Trilux Microbeta counter (Gaithersburg, MD).

B. <u>Results</u>

A representative experiment showing the effects
of purified Zot on proliferation of human PBMC induced by PHA and tetanus toxoid is shown in Figure 3. The results clearly indicate that incubation with Zot markedly suppressed tetanus

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toxoid-induced proliferation (~85% at 60 μ g/ml), while it had no effect on PHA-induced proliferation. Moreover, suppression of TT-induced proliferation by Zot appears to be dose-dependent. Significantly higher levels of suppression were observed when Zot was added at 60 μ g/ml (~85%) than when 20 μ g/ml Zot were used (~56% suppression). Addition of BSA, used as control protein has no effect on either TT or PHA-induced lymphocyte proliferation demonstrating the specificity of Zot biological activity.

To further examine the specificity of Zot-induced suppression of TT-induced proliferation, PBMC were incubated in the absence or presence of TT with ZOT alone, ZOT + anti-ZOT rabbit antiserum (used at a 1:10 dilution) or Zot + normal rabbit serum (used at a 1:10 dilution). As can be observed Figure 4, addition of an anti-Zot antiserum reversed by greater than 50% Zot-mediated suppression of TT-induced proliferation. of normal rabbit serum had no effect, confirming the specificity of Zot-mediated effects. addition of BSA had no effect in this system.

EXAMPLE 3
25 Effects Of Zot On FITC-Dextran Uptake
By Human Monocytes and Macrophages

A. <u>Materials and Methods</u>

Isolation of Human Peripheral Blood Mononuclear

Cells (PBMC) from Healthy Volunteers

PBMC density gradient were isolated by separation centrifugation over lymphocyte media Organon-Teknika, Durham, NC) from healthy In accordance with the institutional volunteers.

review board of University of Maryland, Baltimore, donors were adults and gave informed consent for the blood drawing. PBMC used were fresh or were aliquoted and frozen in RPMI containing 10% (v/v) FCS and 10% (v/v) DMSO using a controlled linear rate freezer apparatus (1°C per min, Planner Biomed, Salisbury, England) to preserve cell viability and maximize cell recovery. Cells were stored in liquid nitrogen until used.

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Preparation of Purified 6xHis-Zot

The 6xHis-Zot fusion protein was prepared and purified by the process described above in Example 2 above.

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Soluble Antigen Uptake

The ability to take up soluble antigen was using fluorescein isothiocyanate measured 50,700; Sigma). (FITC) - conjugated dextran (MW Freshly isolated PBMC (500,000 cells in 0.5 ml of 20 cRPMI containing 10% (v/v) heat-inactivated FCS and incubated in gentamicin) were 50 μg/ml of absence or presence of purified Zot (40 $\mu g/ml$) or BSA (40 μ g/ml) for 3 hr at 37°C in a final volume of This incubation was performed 0.5 ml/50 ml tube. 25 under agitation in siliconized tubes to prevent the the tube monocytes/macrophages to adherence of walls. Following this incubation, FITC-dextran (at a final concentration of 300 µg/ml), as well as allophycocyanin (APC)-labeled and anti-CD14 30 Chlorophyll anti-HLA-DR Peridinin Protein (PerCP) - labeled monoclonal antibodies were added to each culture without washing and cells

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allowed to incubate for 30 min at 37°C or on ice. Since uptake of FITC-dextran, and soluble antigens in general, depends on pinocytosis, a temperature-dependent phenomenon, incubations at 0°C are performed to establish the levels of nonspecific binding of FITC-dextran to the cells.

In these experiments CD14 and HLA-DR (a major histocompatibility complex Class II antigen whose level of expression increase with cell activation) were used to identify monocyte/macrophages.

Cells were then washed once with ice cold PBS and run immediately on a Coulter Epics Elite flow cytometer/cell sorter system (Coulter Corp., Miami, FL). Analysis was performed using the WinList software package (Verity Software House, Topham, ME). The percentages of cells that incorporated FITC-dextran were obtained by subtracting the percentage of cells that incorporated FITC-dextran at 0°C (ice) from the percentage of cells that incorporated FITC-dextran at 37°C.

B. Results

A representative experiment showing the effects of Zot on FITC-dextran uptake by normal human CD14+ 25 HLA-DR+ monocyte/macrophages is shown in Figures 5A-5D. The results show that Zot (Figure 5D) markedly suppressed (~51-58%) FITC-dextran uptake by human monocyte/macrophages as compared to that observed in cells incubated with media alone or BSA. No significant differences were 30 observed between the percentage of cells that incorporated FITC-dextran in media orin the presence of BSA. Moreover, as expected, incubation

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totally abrogated FITC-dextran uptake, 0°C that this phenomenon confirming temperature-dependent.

It is well established that antigen uptake by APCs, such as monocyte/macrophages, is a critical lymphocyte activation leading to event proliferation (Sztein et al, supra (1997)). The Zot interferes with showing that results the uptake demonstrate that FITC-dextran of Zot on TT-induced immunoregulatory effects proliferation are mediated, at least in part, decreasing the ability of monocyte/macrophages to uptake antigen, leading to alterations in antigen and presentation. This further processing supported by the fact that Zot does not affect PHA-induced proliferation, a phenomenon that does not require antigen processing and presentation.

EXAMPLE 4

Measurements of the Number of FITC-Zot 20 Binding Sites/Cell in Human Monocytes/Macrophages and Lymphocytes

Α. Material and Methods

25 Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

by density gradient isolated PBMC were centrifugation over lymphocyte separation media (LSM, Organon-Teknika, NC) from healthy Durham, In accordance with the institutional volunteers. review board of University of Maryland, Baltimore, donors were adults and gave informed consent for the PBMC were aliquoted and frozen in blood drawing. RPMI containing 10% (v/v) FCS and 10% (v/v) DMSO 35

using a controlled linear rate freezer apparatus (1°C per min, Planer Biomed, Salisbury, England) to preserve cell viability and maximize cell recovery. Cells were stored in liquid nitrogen until used. In some experiments, cells were used immediately after isolation.

Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described 10 in Example 2 above.

Conjugation of Zot to Fluorescein Isothiocyanate (Zot-FITC)

Conjugation of Zot with FITC was performed 15 Briefly, Zot was following standard techniques. labeling buffer FITC 500 ml dialyzed against comprising 0.05 M boric acid, 0.2 M NaCl, adjusted to pH 9.2 with concentrated NaOH, and stored at 4°C, at 4°C overnight to remove free $\mathrm{NH_4}^+$ ions and raise 20 the pH to 9.2. 20 μl of 5.0 mg/ml FITC in DMSO for each milligram of Zot was then added, followed by an incubation for 2 hr at room temperature. Unbound FITC was then removed by dialysis in 500 ml dialysis 7.4), Tris-HCl Hq) М comprising 0.1 buffer 25 0.1% (w/v) NaN₃, 0.2 M NaCl, adjusted to pH to 7.4 with concentrated NaOH, and stored at 4°C, at 4°C with two to three changes over 2 days. This preparation was stored at 4°C until used.

Binding of FITC-Zot to Human PBMC and Flow Cytometric Analysis

PBMC isolated as described above were incubated with increasing concentrations of Zot-FITC 5 30 min at 37°C in siliconized tubes (to preclude the binding of macrophages to the test tube walls) in the presence of monoclonal antibodies (mAb) to CD14 phycoerythrin (PE) and to CD3 conjugated to conjugated to ECD (energy coupled dye, 10 PE-Texas-Red conjugate). CD14 is used as a marker of human macrophages while CD3 is used as a marker for T lymphocytes. The use of these fluorochromes (e.g., FITC, PE and ECD) allowed us to study simultaneously the binding of Zot to macrophages. 15 T lymphocytes in mixed PBMC populations 3-color flow cytometry. Following staining, cells (pH 7.2) containing were washed twice with PBS 1.0% (w/v) BSA and 0.1% (w/v) NaAzide and analyzed immediately by flow cytometry using an Epics Elite 20 flow cytometer/cell sorter system (Beckman-Coulter, experiments, these Miami, FL). In fluorochrome-labeled mAbs of the same isotypes, but irrelevant specificity, were used as a controls. Platelets, erythrocytes (if any) and cell debris 25 analysis by setting excluded from appropriate gate on the forward vs. 90% For each sample we collected scatter parameters. data for over 10,000 cells. Data analysis was performed using the Epics Elite analysis package 30 (Coulter) or the WinList list-mode analysis package (Verity Software House, Topsham, ME). The amounts (in pM) of Zot-FITC added to each tube was derived

from the final concentrations added (in $\mu g/ml$) and the known MW of Zot (44,900).

Calculation of Zot Binding Sites/Cell

of intensity fluorescence each mean 5 population following incubation with Zot-FITC were converted to number of Zot binding sites/cell using a standard curve constructed using the Quantum 26 MESF kit (range 10,000 to 500,000 MESF) and the OuickCal calibration software according to the 10 Cytometry (Flow recommendations manufacturers Standards Corporation, San Juan, Puerto Rico). The fluorescent standards in the kit are Ouantum Soluble of Eguivalent Molecules calibrated in against solutions Fluorochrome (MESF) units 15 The number of binding purified fluorescent dyes. sites per cell (macrophages or lymphocytes) units of Zot-FITC incubated derived from MESF samples (with non-specific binding, i.e., MESF of subtracted) controls, FITC-labeled mouse IqG 20 adjusted for the fluorescein/protein ratio (F/P) of the various FITC-Zot batches used.

B. Results

A representative experiment showing the flow 25 cytometric analysis of Zot-F ITC binding to human T lymphocytes (CD3⁺) and macrophages (CD14⁺) is shown The results show that binding of in Figure 6. Zot-FITC, as evidenced by increased number of Zot higher fold is several binding sites/cell, 30 lymphocytes. CD3⁺ Т than in macrophages. Furthermore, these data show that binding of Zot is a saturable phenomenon, with saturation reached at

approximately 0.5 pM (~40 g/ml). Moreover, we observed that under saturation conditions, there are ~106,000 Zot-binding sites/cell and ~9,000 Zot-binding sites/cell in human macrophages and lymphocytes, respectively.

In order to explore the distribution of Zot binding sites in human macrophages and lymphocytes, the methodology described above was used to determine the number of Zot binding sites/cell in several volunteers. The results indicated that the average number of Zot binding sites/cell is, on average, approximately 10-fold higher in macrophages (mean=104,649; range=56,791-142,840) than in lymphocytes (mean=10,684; range=4,802-18,662).

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EXAMPLE 5 Kinetics of Zot Binding to Human Monocytes/Macrophages and Lymphocytes

20 A. <u>Material and Methods</u>

Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described above in Example 4.

Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described in Example 2 above.

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Conjugation of Zot to fluorescein isothiocyanate (Zot-FITC)

Conjugation of Zot with FITC was as performed in Example 4 above.

Kinetics of Binding Assays

Determination of the kinetics of binding of Zot to human cells was carried out by flow cytometry Zot-FITC conjugates and flow cytometry. using Samples were run in real time using an EPIC ELITE 5 flow cytometry/cell sorter system (Beckman-Coulter, PBMC were stained with anti-CD3 mAb Miami, FL). tagged with ECD (energy-coupled dye) and anti CD-14 mAb tagged with PE (phycoerythrin). Controls for fluorescence were prepared 10 background cell suspension additional aliquot of each substituting irrelevant mAb of the same isotypes, conjugated with the corresponding fluorescent dyes, experimental mAb. PBMC labeled for the anti-CD3 and anti-CD14 mAbs were then washed and 15 maintained in an ice bath until analyzed (within antigen modulation. Before 1 hr) to minimize analysis, cells were allowed to equilibrate at 37°C for 15-20 min in a water bath and maintained at 37°C using a viable sample handler (kinetics module, 20 Cytek, Fremont, CA) attached to the flow cytometer for the duration of the experiment (12 min) while data was continuously collected. Baseline FITC fluorescence levels were collected for 25 90-150 seconds and data acquisition was paused for final ~10-15 sec to inject Zot-FITC (to concentration of 40 µg/ml). Data collection was resumed immediately after the addition of Zot-FITC at a rate of ~300-600 cells/sec for a total of 30 12 min. FITC, PE and ECD-fluorochrome were excited using an air-cooled argon laser (488 nm emission). Results were calculated and displayed using WinList and Isocontour analysis packages (Verity

Software House, Topsham, ME). Data are presented as isometric displays of Zot-FITC intensity (y axis) versus time (x axis) versus cell number (z axis) for cells gated on CD3 (T lymphocytes) or CD14 (macrophages).

B. Results

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experiment showing representative the kinetics of Zot-F ITC binding to human T lymphocytes (CD14⁺) is shown 10 and macrophages The results indicate that Figures 7A-7C. Zot 7A) binding to human macrophages (Figure and lymphocytes (Figure 7B) occurs very reaching equilibrium within 2 min following addition 15 of Zot-FITC. To compare the time required for Zot and an anti-CD14 mAb to reach maximum binding, similar experiments were performed using unlabeled cells. In these experiments, baseline FITC fluorescence levels were collected as described above, data acquisition was paused for ~10-15 sec to 20 FITC-anti-CD14 mAb and data collection resumed immediately for a total of 12 min. results (Figure 7C) indicate that maximum levels of Zot binding occur in less time (~2 min) than that 25 required by anti-CD14 mAb to reach equilibrium (~5 min).

EXAMPLE 6 Binding of FITC-Zot to Human B and T lymphocytes

A. Material and Methods

Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described in Example 4 above.

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Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described in Example 2 above.

Conjugation of Zot to fluorescein isothiocyanate (Zot-FITC)

Conjugation of Zot with FITC was as performed in Example 4 above.

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Binding of FITC-Zot to Human PBMC and Flow Cytometric Analysis

PBMC isolated as described above were incubated with 40 μ g/ml of Zot-FITC for 30 min at 37°C in the 25 presence of mAb to CD14 conjugated to PE, conjugated to Tricolor (a PE-Cy5 conjugate) and anti-CD19 conjugated to ECD (energy coupled dye, a is a marker of PE-Texas-Red conjugate). CD14 human macrophages, CD3 is a marker of T lymphocytes 30 and CD19 is a marker for B lymphocytes. The use of these fluorochromes (e.g., FITC, PE, TC and ECD) allowed us to study simultaneously the binding of Zot to macrophages, T and B lymphocytes in mixed by 4-color flow cytometry. 35 PBMC populations Following staining cells were washed twice with PBS

(pH 7.2) containing 1.0% (w/v) BSA and 0.1% (w/v) NaAzide and analyzed immediately by flow cytometry using an Epics Elite flow cytometer/cell sorter system (Beckman-Coulter, Miami, FL). In these experiments, fluorochrome-labeled mAbs of the same isotypes, but irrelevant specificity, were used as a controls. Platelets, erythrocytes (if any) and cell debris were excluded from analysis by setting an appropriate gate on the forward vs. 90% light scatter parameters. For each sample, data for over 10 Data analysis was 10,000 cells was collected. performed using the Epics Elite analysis package (Coulter) or the WinList list-mode analysis package (Verity Software House, Topsham, ME). The results are shown as single color histograms of Zot-FITC 15 fluorescence intensity in T (CD3⁺) and B (CD19⁺) lymphocyte-gated populations.

B. Results

20 An experiment showing the binding of Zot-F ITC to human T (CD3⁺) and B (CD19⁺) lymphocytes is shown in Figure 8. The results indicate that Zot binds similarly to both, T and B lymphocytes. In this experiment Zot-FITC binding to macrophages was 25 5-8 fold higher than binding to T or B lymphocytes.

EXAMPLE 7 Inability of Zot Antagonists to Block Zot-FITC Binding

5 A. Material and Methods

Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described above in Example 4.

Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described in Example 2 above.

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Preparation of FZI/0 and FZI/1 Zot Antagonists

Peptide antagonists FZI/0 (Gly Gly Val Leu Val Gln Pro Gly) (SEQ ID NO:7) and FZI/1 (Val Gly Val Leu Gly Arg Pro Gly) (SEQ ID NO:8) were chemically synthesized and purified using well-known techniques, such as described in High Performance Liquid Chromatography of Peptides and Proteins: Separation Analysis and Conformation, Eds. Mant et al, C.R.C. Press (1991), and a peptide synthesizer, such as Symphony (Protein Technologies, Inc).

Conjugation of Zot to Fluorescein isothiocyanate (Zot-FITC)

Conjugation of Zot with FITC was as performed in Example 6 above.

Culture Conditions for Blocking of FITC-Zot Binding to Human PBMC and Flow Cytometric Analysis

5 PBMC isolated as described above were stained with mAbs to CD14 conjugated to PE and ECD (energy coupled dye, conjugated to PE-Texas-Red conjugate). Cells were then washed and incubated for 15 min at $4^{\circ}C$ in 400 μl of AIM-V medium 10 (GIBCO BRL, a defined serum-free medium routinely for human lymphocyte cultures) 0.2% (w/v)NaAzide (to internalization/recycling) in media alone or with addition of FZI/0 (4.0)mg/ml), (4.0 mg/ml), BSA (4.0 mg/ml; negative control) or 15 unlabeled Zot (160 μg/ml; positive control). Zot-FITC was then added to each tube to reach a final concentration of 40 μ g/ml and incubated for 5 min (the time required to reach equilibrium), 20 washed and immediately run in the flow cytometer. Thus, FZI/0, FZI/1 and BSA were added at 100-fold excess and unlabeled Zot at 4-fold excess compared the concentration of Zot-FITC added. to Unfortunately, the technical difficulties 25 obtaining large amounts of purified unlabeled Zot preparations precluded evaluating its ability to block Zot-FITC binding at more than 4-fold excess. Platelets, erythrocytes (if any) and cell debris excluded were from analysis by setting 30 appropriate gate on the forward vs. 90% light scatter parameters. For each sample we collected data for over 10,000 cells. Data analysis was performed using the Epics Elite analysis package (Coulter) or the WinList list-mode analysis package

(Verity Software House, Topsham, ME). The results are shown as % suppression of the mean fluorescence intensity of cells incubated with Zot-FITC in the presence of Zot antagonists, unlabeled Zot or BSA as related to the mean fluorescence intensity of cells incubated in media alone (arbritarily assigned a value of 100%).

B. Results

A representative experiment showing the effects 10 of incubation of PBMC from 3 different volunteers with Zot antagonists, unlabeled Zot or BSA is shown The addition of either FZI/0 or FZI/1 in Figure 9. did 100-fold excess antagonists at significantly block binding of Zot-FITC to CD14+ 15 gated macrophages. Similarly, addition of 100 fold excess BSA did not affect binding of Zot-FITC. contrast, pre-incubation with only 4-fold excess unlabeled Zot blocked binding of Zot-FITC by 24-43%. These results suggest that binding of Zot to human 20 macrophages involves a receptor with different binding sites that those of Zot/zonulin receptors identified in brain and intestinal tissues.

EXAMPLE 8

Zot Suppression of Tetanus Toxoid

(TT)-Induced Proliferation is

Dependent on Factor(s) Present in Serum

30 A. Material and Methods

Isolation of Human Peripheral Blood Mononuclear cells (PBMC) from healthy volunteers

PBMC were isolated from healthy volunteers as described in Example 4 above.

Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described in Example 2 above.

5 Culture Conditions and Lymphoproliferation Assays

PBMC $(1.5 \times 10^6 \text{ cells/ml})$ were cultured in 1.0 ml of either (a) AIM-V medium, (b) RPMI 1640 containing 10% (v/v) heat-inactivated fetal calf 10 serum and 50 μg/ml gentamicin or (c) RPMI 1640 containing 10% (v/v) heat-inactivated human AB serum and 50 μ g/ml gentamicin. Cells were incubated at 37° C, 5% CO₂ in 96-well plates in the absence or presence of tetanus toxoid (TT; a specific antigen, 15 used at 2.0 μg/ml; Wyeth, Marietta, PA) without or with purified Zot (60 $\mu g/ml$) or bovine serum albumin (BSA; a control protein, 60 μg/ml; Fraction V, Sigma, St. Louis, MO). Cells were cultured for 6 days and 1.0 Ci/well of tritiated thymidine was 20 Plates were harvested 20 hr later on a added. Wallac cell harvester (Gaithersburg, MD) and incorporated thymidine measured on a Wallac Trilux Microbeta counter (Gaithersburg, MD).

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B. Results

The results from 3 independent experiments showed that no inhibition of TT-induced proliferation could be observed when PBMC were incubated in the absence of serum, e.g., when the AIM-V defined media was used in the cultures. fact the reverse was observed in most cases, i.e., incubation with Zot in the absence of serum lead to increased proliferative responses TT. In

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contrast, incubation with Zot resulted in marked suppression of TT-induced proliferation when cultures were performed in the presence of either FCS or human AB serum. In fact, the presence of human AB serum appears to mediate higher levels (up to 90%) of suppression of TT-induced proliferation than that observed in the presence of FCS.

EXAMPLE 9 Zot Suppression of CD14 Expression on Human Monocytes/Macrophages

A. <u>Material and Methods</u>

Isolation of Human Peripheral Blood Mononuclear Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described in Example 4 above.

20 Preparation of Purified 6xHis-Zot

The preparation of 6xHis-Zot was as described in Example 2 above.

Culture Conditions and Flow Cytometric Analysis

PBMC isolated as described above were incubated at 37°C, 5% CO₂ in 24-well plates for various time periods (4 hr to 7 days) in the absence or presence of tetanus toxoid (TT; a specific antigen, used at 2.0 μg/ml; Wyeth, Marietta, PA) without or with purified Zot (60 μg/ml) or bovine serum albumin (BSA; a control protein, 60 μg/ml; Fraction V, Sigma, St. Louis, MO). Cells were then stained with a mAb to CD14 conjugated to FITC and analyzed by

flow cytometry. Platelets, erythrocytes (if any) and cell debris were excluded from analysis by setting an appropriate gate on the forward vs. 90% light scatter parameters. For each sample, data for Data analysis was over 10,000 cells was collected. performed using the Epics Elite analysis package (Coulter) or the WinList list-mode analysis package The results (Verity Software House, Topsham, ME). shown as single color histograms of fluorescence on cells gated on the region", defined based on the forward scatter vs. side scatter characteristics of human macrophages.

B. Results

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A representative experiment showing the effects 15 of incubation of PBMC with Zot on the expression of included is CD14 in human macrophages The addition of Zot caused a Figures 10A-10B. expression of suppression of the marked following 18 hr of incubation. This effect was very 20 pronounced either in the absence (Figure 10A) or presence (Figure 10B) of tetanus toxoid. Addition Kinetic of BSA did not affect CD14 expression. experiments showed that Zot-induced suppression of CD14 expression is observed in approximately half of 25 following 4-6 hr experiments as early as exposure to Zot and that CD14 expression remains markedly suppressed after 7 days in culture (the last time point evaluated). CD14 is a molecule in of macrophages that acts 30 surface high-affinity receptor for LPS-LPS-binding protein complexes. Thus, down-regulation of CD14 expression by Zot is believed to have profound effects on

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macrophage activation and its ability to effectively initiate T cell-mediated immune responses.

EXAMPLE 10

Effects of Zot on the Viability of Human Lymphocytes and Monocytes/Macrophages

A. Material and Methods

Isolation of Human Peripheral Blood Mononuclear

Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described in Example 4 above.

15 <u>Preparation of Purified 6xHis-Zot</u>

The preparation of 6xHis-Zot was as described in Example 2 above.

Culture Conditions and Flow Cytometric Analysis

- PBMC isolated as described above were incubated 20 at 37°C , 5° CO_2 in 24-well plates for various time periods (6 hr to 7 days) in the absence or presence of tetanus toxoid (TT; a specific antigen, used at 2.0 μ g/ml; Wyeth, Marietta, PA) without or with purified Zot (40 μ g/ml) or bovine serum albumin 25 (BSA; a control protein, 40 μ g/ml; Fraction V, Sigma, St. Louis, MO). Cells were then stained with dialyzed mAbs to CD14 conjugated to FITC and washed. To assess cell viability, propidium iodide (PI; 50 g/ml; a dye that is readily incorporated into dead cells but excluded from viable cells) was added to the cell suspensions and the samples analyzed Platelets, cytometry. flow immediately by erythrocytes (if any) and cell debris were excluded
 - 35 from analysis by setting an appropriate gate on the

forward vs. 90% light scatter parameters. For each sample, data for over 10,000 cells was collected. Data analysis was performed using the Epics Elite analysis package (Coulter) or the WinList list-mode analysis package (Verity Software House, The results are shown as the % viable cells "monocyte region" or "lymphocyte on the gated region", defined based on the forward scatter vs. characteristics of these scatter side populations.

B. Results

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A representative experiment showing the effects of incubation of PBMC with Zot on the viability of human macrophages and lymphocytes is included in 15 moderate Zot caused Figure 11. Addition of viability of macrophages in the decreases compared to controls as early as 1 day in culture. These decreases became quite pronounced by day 4 and virtually all macrophages were dead after 7 days in 20 In contrast, no differences in lymphocyte culture. viability were observed among cultures incubated with Zot and those incubated with media or BSA until However, lymphocyte viability was markedly decreased at later times in the presence of Zot. 25 Similar results were observed when TT was added to the cultures in the absence or presence of Zot. These results demonstrate that Zot affect macrophage viability at relatively early times, while the effects on lymphocytes do not become apparent until 30 These observations at least 4 days in culture. concerning the additional information provide that may underlie Zot-mediated mechanisms

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suppression of antigen processing and presentation, pointing to an early effect on macrophages.

EXAMPLE 11

Zot-mediated Induction of Cytokine Production by Human Monocytes/Macrophages and Lymphocytes

A. Material and Methods

Isolation of Human Peripheral Blood Mononuclear

Cells (PBMC) from Healthy Volunteers

PBMC were isolated from healthy volunteers as described in Example 4 above.

15 <u>Preparation of Purified 6xHis-Zot</u>

The preparation of 6xHis-Zot was as described in Example 2 above.

Culture Conditions

- PBMC isolated as described above were incubated 20 at 37°C , 5% CO_2 in 24-well plates for various time periods (6 hr to 4 days) in the absence or presence of tetanus toxoid (TT; a specific antigen, used at 2.0 μ g/ml; Wyeth, Marietta, PA) without or with purified Zot (60 μ g/ml) or bovine serum albumin 25 a control protein, 60 μ g/ml; Fraction V, Sigma, St. Louis, MO). For studies involving the production of TNF- α , IL-1 β and IL-10, the content of the wells were collected at 6 hr and at days 1, 2 and 4 into 1.5 ml Eppendorf tubes and centrifuged at 30 4°C for 10 min at 2,700 x g in a refrigerated Eppendorf centrifuge to remove cells and debris. Supernatants were then transferred to new Eppendorf -70°C until analyzed. at and frozen tubes
 - 35 Supernatants for the measurement of

T lymphocyte-derived cytokines (e.g., IL-2, IL-4 and IFN- γ) following stimulation with TT without or with purified Zot or BSA were collected after 3 days.

5 Cytokine Measurements by Chemiluminescence ELISA

A standard chemiluminescence capture ELISA was used to detect the presence of cytokines in the cell Briefly, $1.0-2.0 \mu g/ml$ of culture supernatants. 10 anti-human cytokine antibodies were coated onto 96-well black opaque ELISA plates (Corning-Costar, Cambridge, MA) in either PBS (pH 7.4) (Biofluids) or 0.1 M sodium bicarbonate (pH 8.1) overnight at 4°C. Plates were washed with PBS (pH 7.4) containing 15 0.5% (v/v) Tween-20 (Sigma) and blocked for 2 hr with either PBS (pH 7.4) containing 10% (v/v) FCS or After washing, 100 μ l of cell 4.0% (w/v) BSA. culture supernatants or recombinant human cytokines (as standards) were added to the wells and incubated 20 for 2 hr at room temperature. After washing, the corresponding anti-cytokine mAbs conjugated biotin were added to the wells (45 min at room incubated with washed and temperature), avidin-peroxidase for 30 min at room temperature. 25 (Boehringer ELISA reagent chemiluminescence was then added Mannheim, Gaithersburg, MD) 1450 Microbeta chemiluminescence detected on a Trilux plate reader (Wallac, Gaithersburg, MD). anti-IL-1ß antibodies were obtained from Endogen 30 from obtained all others were (Woburn. MA), Pharmingen (San Diego, CA).

B. Results

A representative experiment showing the ability induce cytokine production by human macrophages is shown in Figures 12A-12C. Addition of Zot (in the absence of TT) induced the production of considerable amounts of TNF- α as early as 6 hr, reaching peak levels at 24 hr (3,400-3,800 pg/ml) and decreasing afterwards reaching near-baseline Lower levels of levels by 4 days (Figure 12A). observed in media BSA cultures 10 $TNF-\alpha$ were or(-1,000-1,100 pg/ml), probably the result non-specific activation of macrophages following adherence to plastic. A weak induction of IL-1 β by Zot (~40~60 pg/ml) was also observed that reached peak levels after 2 days and decreased considerably 15 by day 4 (~20 pg/ml) (Figure 12B). No significant levels of IL-1ß were observed in media or BSA Finally, incubation with Zot resulted in cultures. the production of high levels of IL-10 after 6 hr 20 (~1,000 pg/ml), reaching peak levels by day 1 (~1,500 remaining the same pg/ml) and at concentrations up to day 4 (Figure 12C). Similar to the findings described above for TNF- α , considerably lower levels of IL-10 were observed in media or BSA probably the result of non-specific 25 cultures, activation of macrophages following adherence to plastic. The presence of TT did not alter either the kinetics or the magnitude of the cytokine responses observed following the addition of Zot. 30 The induction of high levels of proinflammatory cytokines following exposure to Zot provides additional information concerning Zot-mediated mechanisms that underlie may

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suppression of antigen processing and presentation, again pointing to an early effect on macrophages. induction example, Zot-mediated production, a cytokine known to be a major inhibitor of the Th1 type response through the inhibition of cytokines such as IL-2. is production of believed to play significant role in the а suppression of antigen-induced lymphoproliferative responses observed added the when Zot is to cultures.

A representative experiment showing the effects of Zot on the production of T cell-derived cytokines is shown in Figures 13A-13B. It was observed that Zot suppressed IL-2 production addition of induced by incubation with TT, while BSA had no No measurable levels of IL-2 effect (Figure 13A). were induced by Zot in the absence of TT. contrast, addition of Zot consistently induced the production of low to moderate levels of IFN-γ in the absence of TT, similar to the levels induced by TT (Figure 13B). Moreover, Zot markedly increased the levels of IFN-induced by TT, while BSA had no effect Finally, it as observed that the (Figure 13B). addition of Zot did not induce IL-4 production. Since IL-2 is a cytokine that plays a critical role in lymphocyte proliferation, the suppression of production of IL-2 following antigenic stimulation believed to be one of the key mechanisms underlying Zot-mediated suppression of TT induced Also of importance, the fact that proliferation. induces IFN-y production in the absence presence of TT, as well as the production of many Zot pro-inflammatory cytokines, indicates that

exerts its immunoregulatory effects at various levels during the complex process of antigen processing and presentation leading to the generation of antigen-specific immune responses.

All references cited herein are incorporated by reference in their entirety.

While the invention has been described in detail, and with reference to specific embodiments thereof, it will be apparent to one with ordinary skill in the art that various changes and modifications can be made therein without departing from the spirit and scope thereof.